Photopolymerizable composition, products made therefrom and processes for preparing such products.

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Description

The present invention relates to a flexible, fast processing, photopolymerizable composition and to a sheet or roll prepared from the composition. The present invention also relates to a coated composition comprised of the flexible, fast processing, photopolymerizable composition on a substrate and to a process for making a printed circuit board using the flexible, fast processing, photopolymerizable composition.

Photopolymerizable compositions have long been known in the art. During their evolution, various chemical compounds have been used as a photopolymerizable component of the composition. One development in the art is disclosed in U.S. Patent No. 2,760,863. This patent relates to a photopolymerizable composition useful in the preparation of printing plates which is comprised of an ethylenically unsaturated, photopolymerizable monomer, a photoinitiator, a polymerization inhibitor and a polymeric binding agent. The composition is coated onto a substrate, dried, exposed to actinic radiation through a stencil or negative and developed to remove the unexposed portions of the composition thereby forming the printing plate.

U.S. Patent No. 3,469,982 marks an advance over the photopolymerizable composition described in U.S. Patent No. 2,760,863. Rather than being applied as a liquid, this later patent discloses a photopolymerizable composition which is coated onto a flexible film base or support and then covered with a removable cover film. In use, the cover film is peeled off, the photopolymerizable composition is laminated to a substrate, exposed to actinic radiation through a negative, the film support is peeled off and the composition is developed in an organic solvent. To form a printed circuit board, the developed composition on the substrate is then subjected to etching and finally stripped in another organic solvent.

In U.S. Patent No. 3,036,914, a photopolymerizable composition having improved flexibility which is useful for preparing printing plates is disclosed. The composition is comprised of: 1) an addition polymerizable, ethylenically unsaturated compound, such as diacrylates and acrylic or methacrylic acid esters of polyalkylene glycols; 2) a photoinitiator; 3) a high molecular weight polyethylene oxide; and 4) at least one essentially linear cellulose ester.

Improved flexibility is also the goal of U.S.S.R. Patent No. 190,211. This goal is allegedly achieved by a photopolymerizable composition comprised of a photopolymerizable oligomer, a photoinitiator and a mixture of copolymers; the first being a copolymer of methacrylic acid and methyl methacrylate and the second being a copolymer of methacrylic acid or acrylic acid with butyl acrylate, alkyl methacrylate or alkyl acrylates containing 5 to 8 carbon atoms in the alkyl group.

In U.S. Patent No. 3,887,450, and divisional U.S. Patent No. 3,953,309, a significant advance in the art is set forth. These patents disclose a photopolymerizable composition which can be made into a storable sheet or roll on a film support. The composition may be used to prepare a printed circuit board and yet can be developed and stripped using aqueous solutions containing an alkaline agent. The photopolymerizable composition is comprised of a photopolymerizable ethylenically unsaturated monomer, a photoinitiator, a polymerization inhibitor and a binding agent which is a polymer of one or more of a defined styrene or vinyl monomer with one or more of a defined alpha, beta-ethylenically unsaturated carboxylic acid- or anhydride-containing monomer. The ratio of the monomers is selected to yield the required performance characteristics.

A further advance in the art is disclosed in U.S. Patent No. 4,239,849. This patent describes a flexible photopolymerizable composition that is resistant to cold flow which may be formed into a dry film on a film support and, after exposure to actinic radiation, which may be developed in dilute aqueous alkaline solutions. The composition is comprised of one or more additional photopolymerizable, ethylenically unsaturated compounds, a photoinitiator and a preformed macromolecular polymeric binding agent. The binding agent is a copolymer of a styrene-type monomer, an acrylate-type monomer and an unsaturated carboxyl-containing monomer.

The present invention consists in a photopolymerizable composition capable of being formed into a storable sheet or roll on a peelable support which on a 100 parts by weight basis comprises:

A. from 10 to 60 parts by weight of an addition polymerizable material including one or more non gaseous compounds containing at least two terminal ethylenic groups and having a boiling point above 100°C;

B. from 0.001 to 20 parts by weight of a photo initiated free radical generating addition polymerization initiating system;

C. from 0.001 to 5 parts by weight of a thermal addition polymerization inhibitor; and

D. from 40 to 90 parts by weight of a preformed macromolecular polymeric binding agent which is a polymer of:

1) a first monomeric material which contains one or more non acidic compounds having the general formula

\[
R - C = CH_2
\]

or

\[
CH_2 = C-Y
\]

X

...
wherein R is hydrogen and alkyl group having from 1 to 6 carbon atoms or a halo group and wherein when
X is hydrogen Y is OOCR, OR, OOCR, COOR, CN, CH = CH₂

or Cl when X is methyl, Y is COOR, CN, CH = CH₂ or

and when X is chlorine, Y is Cl and wherein R₃ is an alkyl group having from 1 to 12 carbon atoms, a phenyl
group or a benzyl group and R₂ and R₄ are hydrogen, an alkyl group having from 1 to 12 carbon atoms or a
benzyl group; and

i) a second monomeric material which consists essentially of one or more alpha, beta ethylenically
unsaturated carboxylic acid- or anhydride-containing monomers having from 3 to 15 carbon atoms, the
ratio of the first monomeric material to the second monomeric material being sufficient to render
substantially all of the binding agent soluble in a substantially wholly aqueous solution containing 1% by
weight of sodium carbonate, characterised in that component A comprises:

1) from 5 to 50 parts by weight of an acrylic having a boiling point greater than 175°C and having the
formula

\[ \text{H₂C} = \text{C–} \text{(OC₉H₂₉)}₉ \text{OR₂} \]

wherein m is from 1 to 4, n is from 1 to 12, R₃ is selected from H, CH₃ and mixtures thereof and R₄ is
selected from unsubstituted phényl, substituted phényl, unsubstituted naphtyl, substituted naphtyl, branched
or unbranched, substituted or unsubstituted alkyl having 1 to 12 carbon atoms, substituted or unsubstituted
cycloalkyl having 5 or 6 carbon atoms in the ring and mixtures thereof and

ii) 5 to 50 parts by weight of said non gaseous compounds containing two or more terminal ethylenic
groups.

The present invention also provides a sheet or roll of the photopolymerizable composition on a film
support, a laminate comprised of the photopolymerizable composition coated onto a substrate and a
process for making a printed circuit board by employing the improved photopolymerizable composition.

As stated hereinabove, the present invention relates to an improved photopolymerizable composition.

The composition is comprised of an addition polymerizable material, a photoinitiated free radical
generating addition polymerization initiating system, a thermal-addition polymerization inhibitor and a
preformed macromolecular polymeric binding agent.

Based on 100 parts by weight of the dried photopolymerizable composition, the addition polymerizable
material is present in an amount ranging from 10 to 60 parts by weight and is comprised of from 5 to 50
parts, preferably from 8 to 25 parts by weight of an acrylic and from 5 to 50 parts, preferably from 8 to 25
parts by weight of one or more non-gaseous compounds containing at least two terminal ethylenic groups
and having a boiling point above 100°C.

In the acrylic of general formula:

\[ \text{H₂C} = \text{C–} \text{(OC₉H₂₉)}₉ \text{OR₂} \]

m is preferably 2 or 3, and most preferably 2, n is preferably from 2 to 8 and most preferably 2 or 3 and R₃ is
preferably H. When R₂ is substituted or unsubstituted alkyl it preferably has 4 to 8 carbon atoms. More
preferably however, R₂ is unsubstituted phenyl, substituted phenyl or a mixture thereof and most
preferably unsubstituted phenyl. The substituents on the phenyl or naphtyl are selected such that they do
not substantially adversely affect the characteristics of the photopolymerizable composition and may be
halogens, especially chlorine, bromine and iodine, alkyl and alkoxy, either group having 1 to 15, preferably
1 to 6 carbon atoms and mixtures thereof.

Acrylates within the scope of the present invention have a low volatility as indicated by a boiling point
greater than 175°C, preferably greater than 200°C and generally possess a molecular weight greater than
240, preferably greater than 280. Additionally, the acrylates are selected such that they do not result in the
degradation of the photopolymerized composition when subjected to etching or plating and do not cause degradation of the aqueous developing or stripping baths.

To a certain extent, the poly-ethylenically unsaturated compound may be substituted with a

The preferred acrylates are substituted and unsubstituted phenoxy(polyethoxymethyl acrylates and substituted and unsubstituted phenoxy(polypropoxy)propyl acrylates with the most preferred acrylates being phenoxy (di, tri, tetra or penta) ethoxymethyl acrylate. Such acrylates are available commercially or may be synthesized by the process of reacting formaldehyde or the appropriate alkylene oxide (e.g., ethylene oxide) with the alcohol form of the R₂ and then esterifying the resulting product with acrylic acid or methacrylic acid in the presence of a suitable acidic catalyst.

The presence of the acrylate monomer in the photopolymerizable composition imparts significant advantages thereto. In particular, the acrylate improves the flexibility and tack of the photopolymerizable composition which improves the handling characteristics of sheets or rolls made of the composition and facilitates lamination to a substrate. Moreover, the acrylate enables the photopolymerized composition to retain a high degree of flexibility. Upon development, the presence of the acrylate in the unexposed portions of the composition facilitates efficient (i.e., rapid and complete) removal from the substrate. Yet, in the photopolymerized portions of the composition, the acrylate improves the efficiency of the stripping operation. The second component of the addition polymerizable material is capable of forming a high polymer by free-radical photoinitiated, chain propagating addition polymerization. Such compounds are disclosed in U.S. Patent Nos. 2,760,863, 3,887,450 and 3,953,309.

Preferably, the compounds are non-gaseous at 20°C and atmospheric pressure, have 2 to 4 or more terminal ethylenic groups and exert a plasticizing action on the polymeric binding agent. Suitable compounds, which may be used alone or in combination, include an alkylene or a polyalkylene glycol diacrylate prepared from alkylene glycols having 2 to 15 carbons or polyalkylene ether glycols of 1 to 10 ether linkages.

Because of their generally more rapid rate of insolubilization on exposure, presumably due to a relatively rapid establishment of a network polymer structure, an outstanding class of low molecular weight addition polymerizable components are those having a plurality of addition polymerizable ethylenic linkages, particularly when present as terminal linkages, and especially those wherein at least one and preferably most of such linkages are conjugated with a doubly bonded carbon, including carbon doubly bonded to carbon and to such heteroatoms as nitrogen, oxygen, and sulfur. Outstanding are such materials wherein the ethylenically unsaturated groups, especially the vinylidene groups, are conjugated with ester or amide structures. The following specific compounds are further illustrative of this class: unsaturated esters of polyols, particularly such esters of the methylene carboxylic acids, e.g., ethylene diacrylate; diethylene glycol diacrylate; glycerol diacrylate; glycerol triacrylate; ethylene dimethacrylate; 1,3-propylene dimethacrylate or diacrylate; 1,2,4-butanetriol trimethacrylate or triacrylate; 1,4-benzenediol dimethacrylate or diacrylate; pentaerythritol methacrylates or acrylates; 1,5-pentanediol dimethacrylate or diacrylate; the bis-acrylates and methacrylates of polyethylene and polypropylene glycols of molecular weight 200–500, such as tripropylene glycol diacrylate; unsaturated amides, particularly those of the methylene carboxylic acids, and especially those of alpha, omega-diamines and oxygen-interrupted omega-diamines, such as methylene bisacrylamide; methylene bismethacrylamide; 1,6-hexamethylene bisacrylamide; diethylenetriamine trimethacrylamide; bis(methacrylamidopropoxy)ethane; beta-methylacrylamidoethyl methacrylate; N-[(beta-hydroxyethylxyloxy)ethyl] acrylamide; vinyl esters such as divinyl succinate, divinyl adipate, divinyl phthalate, divinyl terephthalate, divinyl benzene-1,3-disulfonate, and divinyl butane-1,4-disulfonate; and unsaturated aldehydes, such as sorbaldehyde (hexadienal).
Such compounds include the substituted or unsubstituted polynuclear quinones, such as 9,10-anthraquinone; 1-chloroanthraquinone; 2-chloroanthraquinone; 2-methylanthraquinone; 2-ethylanthaquinone; 2-tert-butylanthaquinone; octamethylanthaquinone; 1,4-napthoquinone; 9,10-phenantheraquinone; 1,2-benzanthraquinone; 2,3-benzanthraquinone; 2-methyl-1,4-naphthoquinone; 2,3-dichloronaphthaquinone; 1,4-dimethylanthaquinone; 2,3-dimethylanthaquinone; 2-phenylanthaquinone; 2,3-diphenylanthaquinone; sodium salt of anthraquinone alphasulfonic acid; 3-chloro-2-methylanthraquinone; retenequinone; 7,8,9,10-tetrahydronaphthacenequinone; 1,2,3,4-tetrahydrobenz(a)anthracene-7,12-dione.

Also suitable as photoinitiators are the xanthones, thioxanthones, chlorothioxanthones, alkylated thioxanthones, alkyl p-dimethylaminobenzoates and other p-dialkyl aminobenzoate esters, and phenyl dialkoxycacetophenones.

The following photoinitiators, described in previously mentioned U.S. Patent No. 2,760,863, some of which may be thermally active at temperatures as low as 85°C, are also useful; vicinal ketaldonyl compounds, such as diacetyl and benzil; aliphaketalldonyl alcohols, such as benzoin and pivaloin; acyloin ethers, e.g., benzoin methyl and ethyl ethers; alpha-hydrocarbon substituted aromatic acyloins; alpha-methylbenzoin; alpha-allylbenzoin; and alpha-phenylbenzoin.

Silver persulfate is also useful as a free-radical generating initiator activatable by actinic radiation. Certain aromatic ketones, e.g., benzophenone and 4,4'-bisdialkyaminobenzophenones, are also useful.

The thermal-addition polymerization inhibitor is present in an amount ranging from 0.001 to 5 parts by weight. Inhibitors useful in the present invention include p-methoxyphenol, hydroquinone, and alkyl and arylsubstituted phenols, hydroquinones and quinones, tert-butyl catechol, pyrogallol, copper resinate, naphtylnamines, betanaphthol, cuprous chloride, 2,6-di-tert-butyl p-cresol, 2,2-methylenebis(4-ethyl-6-t-butylphenol), phenothiazine, pyridine, nitrobenzene, dinitrobenzene, p-toluquinone, chloranil, aryl phosphites, and aryl alkyl phosphites.

The polymeric binding agent is present in an amount ranging from 40 to 90 parts by weight and is a copolymer of one or more non-acidic compounds with one or more acidic compounds. The non-acidic compounds have the general formula

\[ R - \overset{\text{C}}{=\text{CH}_2} \]  

or

\[ \text{CH}_2 - \overset{\text{C}}{=\text{Y}} \]

wherein R is hydrogen or an alkyl group having from 1 to 6 carbon atoms or a halo group. The benzene ring may be ring substituted with functional groups, such as nitro, alkoxy, acyl, carboxyl, sulpho, hydroxy or halo and the formula should be understood to include such substituents unless otherwise stated. From 1 to 5 benzene substituents may be present, preferably, the substituents are a single alkyl group such as a methyl or t-butyl group. Most preferred of these compounds are styrene, alpha-methyl styrene, para-methyl styrene and para-t-butyl styrene.

The presence of the styrene-type monomer in the polymeric binding agent imparts to the composition improved resistance to the developing and etching or plating solutions. Therefore, depending on the developing and surface modifying conditions selected, the presence and amount of the styrene-type constituent may be selected to yield acceptable development and acceptable resistance to the surface modifying conditions.

With regard to the second general formula, when X is hydrogen, Y is OOCCR\_1, OR\_1, OCR\_1, COOR\_1, CN, CH = CH\_2,

\[ \overset{\text{O}}{\text{CNR}}_1R_4 \]

or Cl; when X is methyl, Y is COOR\_1, CN, CH = CH\_2, or

\[ \overset{\text{O}}{\text{CNR}}_1R_4 \]

and when X is chlorine, Y is Cl; and wherein R\_1 is an alkyl group having from 1 to 12 carbon atoms, a phenyl or a benzyl group and R\_2 and R\_4 are hydrogen, an alkyl group having 1 to 12 carbon atoms or a benzyl group.

Examples of these vinyl-type monomers are vinyl acetate, vinyl butyrate, vinyl benzoate, vinyl chloride, vinylidene chloride, methyl methacrylate and methyl acrylate, acrylonitrile and methacrylonitrile, methacrylamide, and alkyl substituted acrylamides, vinyl methyl ketone, vinyl propyl ketone, vinyl methyl ether, vinyl ethyl ether and vinyl hexyl ether.
The vinyl-type monomer may also be an acrylate-type monomer which includes alkyl and hydroxalkyl acrylates and methacrylates wherein the alkyl group has from 1 to 12, preferably from 1 to 6, carbon atoms. Examples of these materials include methyl methacrylate, ethyl acrylate, butyl acrylate, hydroxypropyl methacrylate, hydroxyethyl methacrylate and hydroxyethyl acrylate. Also, advantageously, mixtures of two or more of these compounds may be used.

The presence of the acrylate-type monomer in the polymeric binding agent imparts to the photopolymerizable composition improved cold flow and flexibility. These features help enable the dry film to be stored as sheets or rolls for long periods of time substantially without fear of degradation caused by cold flow or loss of flexibility.

The acidic monomer may be one or more alpha, beta-ethylenically unsaturated carboxyl containing monomers having from 3 to 15 carbon atoms, preferably from 3 to 6 carbon atoms. Most preferred compounds are acrylic acid and methacrylic acid. Other acids which may be used are cinnamic acid, crotonic acid, sorbic acid, itaconic acid, propionic acid, maleic acid, and fumaric acid, or the corresponding half esters or, where possible, the corresponding anhydrides.

The ratio of the styrene-type or vinyl-type monomer to the acidic comonomer is selected so that the copolymer is soluble in the aqueous alkali medium selected for development. If the amount of the styrene-type or vinyl-type monomer is too high, the unexposed portion of the composition will not be sufficiently soluble; on the other hand, if the amount of styrene-type or vinyl-type monomer is too low, the exposed area of the composition will be tacky, swollen, or dissolved in the aqueous alkali. As one convenient criteria, the binder copolymer should be such that a 40 percent solution in ketones or alcohols will have a viscosity of from 0.1 to 50 Pas. A further convenient criteria is that substantially all of the copolymer will be solubilized in a dilute substantially wholly aqueous solution containing 2% by weight of sodium carbonate at 25°C in about one hour. In this criteria, the term substantially has been used to indicate the level of acceptable development of the photopolymerizable composition and to reflect the fact that while some organic material may be present in the solution, such as by design, accident or reuse of the developing solution, the presence of such organic material is not required for acceptable development.

Representative comonomer weight ratios are from 70:30 to 85:15 for styrene-acrylic acid or methacrylic acid; from 70:30 to 85:15 for methyl methacrylate or combinations of alkyl acrylates or alkyl methacrylates-acrylic acid and/or methacrylic acid; from 35:65 to 70:30 for styrene-monomobutyl maleate and from 70:30 to 95:5 for vinyl acetate-crotonic acid. The degree of polymerization of the binder-copolymer is such that the binding agent forms a non-tacky continuous film upon casting from an appropriate solvent and after appropriate drying. Broadly, the molecular weight is from 1,000 to 500,000. The ranges for the copolymer ratios and the degree of polymerization required for the useful particular binders can be readily ascertained by testing the solubility in a dilute alkali solution. In this regard, mixtures of the monomeric materials as well as mixtures of copolymers may be used to form the binding agent to obtain the desired properties thereof.

In the event that improved resistance to developing and surface modifying solutions as well as improved flexibility and improved cold flow are desired, the polymeric binding agent may be formulated in accordance with the general teachings of previously mentioned U.S. Patent No. 4,239,849. As fully explained therein, the polymer binding agent is a copolymer of: 1) the styrene-type monomer; 2) the acrylate-type monomer; and 3) the acidic monomer within the definitions previously given.

The amounts of the components forming the terpolymer binding agent are selected to at least fulfill the criteria noted previously. As a guideline, the following weight proportions may be followed to prepare the polymeric binding agent.

<table>
<thead>
<tr>
<th>Binder Component</th>
<th>Broad Range</th>
<th>Preferred Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Styrene-type</td>
<td>10—60%</td>
<td>19—35%</td>
</tr>
<tr>
<td>Acrylate-type</td>
<td>10—75%</td>
<td>40—75%</td>
</tr>
<tr>
<td>Carboxylic acid-type</td>
<td>15—40%</td>
<td>18—30%</td>
</tr>
</tbody>
</table>

If desired, the photopolymerizable composition of the present invention may also contain plasticizers, adhesion promoters, dyes and pigments. Suitable colorants will be compatible with the photosensitive compositions and not interfere appreciably with the photosensitivity of the composition. The following specific compounds are illustrative: Fuchsine (C.I. 42510); Auramine Base (C.I. 41008); Caloccid Green S (C.I. 44090); Para Magenta (C.I. 42500); Tryparosan (C.I. 42505); New Magenta (C.I. 42520); Acid Violet RRH (C.I. 42425); Red Violet 5RS (C.I. 42690); Nile Blue 2B (C.I. 51185); New Methylene Blue GG (C.I. 51195); C.I. Basic Blue 28 (C.I. 42585); Iodone Green (C.I. 42556); Night Green B (C.I. 42115); C.I. Direct Yellow 9 (C.I. 19540); C.I. Acid Yellow 17 (C.I. 19865); Acid Yellow 20 (C.I. 18900); Tartrazine (C.I. 19140); Supramine Yellow G (C.I. 19300); Buffalo Black 10B (C.I. 27790); Naphthalene Black 12R (C.I. 20350); Fast Black L (C.I. 51215); Ethyl Violet (C.I. 42600); Pontacyl Wool Blue BL (C.I. 50315); Pontacyl Wool Blue GL (C.I. 42320). (Numbers obtained from the second edition of the Color Index.)
The photopolymerizable composition of the present invention exhibits remarkable versatility. While the composition may be used to prepare printing plates, it possesses the more demanding characteristics required for a successful photoresist in the preparation of a printed circuit board. In particular, after a printing plate composition is coated onto the substrate, it is exposed to actinic radiation through a stencil and contacted with the developing solution to remove those portions of the composition which have been shielded from the radiation. The exposed portions of the composition are then inked and used for producing printed images.

In significant contrast, a photopolymerizable composition which is to be used as a photoresist must of course be capable of being exposed and developed, but the exposed portions of the composition must also be unaffected by etching solutions, such as a 45° Baume solution of ferric chloride, and/or electroplating solutions, such as acid copper sulfate. Despite its resistance to the aggressive solutions used in etching or plating, the photopolymerized composition must be capable of being stripped from the substrate so that the processing of the printed circuit board can be completed.

The photopolymerizable composition of the present invention meets all the stated criteria and hence may be successfully employed as a photoresist in the preparation of printed circuit boards. In this regard, the photopolymerizable composition can be coated onto a flexible film support and be formed into sheets or rolls which are capable of being stored for an extended time before being used. When in this form, the photopolymerizable composition of the present invention is characterized by a uniform thickness in the range of from 12.7 to 127 μm (0.0005 to 0.005 inches) and is free of defects such as voids, pinholes, bubbles, patterns or the like.

The flexible film support may be any material which adequately supports the photopolymerizable composition, does not react with the composition and can be readily separated from the composition, such as by peeling, either before or after exposure to actinic radiation. Materials which satisfy these criteria are high polymers, such as polyamides, polyolefins, polyesters, vinyl polymers and cellulose esters having a thickness in the range of from 12.7—60.8 μm (0.0005 to 0.002 inches). One preferred film support is a film of polyethylene terephthalate having a thickness of about 25.4 μm (0.001 inch).

When in the form of a sheet or roll, the photopolymerizable composition may be protected by a cover sheet on the side opposite to the film support, thereby forming a sandwich structure with the photopolymerizable composition in the middle. The cover sheet may be the same or a different material than the film support and may have a thickness in the same range given previously for the film support. One particularly suitable material is a film of polyethylene having a thickness of about 25.4 μm (0.001 inch).

In use, at least a portion of the sheet or roll is laminated to a substrate. The substrate may be plates, sheets or foils of plastic, steel, aluminum, copper, zinc, magnesium, etc. or may be composites containing such materials. Preferably, the substrate is a copper clad laminate, such as a copper clad, epoxy-fibreglass board, which is a common precursor for a printed circuit board. Lamination may be achieved by removing the cover sheet, if present, and pressing the photopolymerizable composition directly onto one side or both sides (if a two sided board is desired) of the substrate at an elevated temperature in the range of from about 65 to about 150°C. One way in which the lamination step may be accomplished is by passing the substrate in contact with the composition between heated rubber coated rollers.

After lamination, the film support may be removed or, more preferably, if a film support which permits sufficient passage of actinic radiation is employed, it may remain on the laminate during the exposure step. To obtain the desired pattern, a process transparency, e.g., a process negative or positive, stencil, mask, continuous tone, negative or positive image, is laid over the photopolymerizable composition of the laminate. The purpose of the process transparency (artwork) is to prevent exposure to actinic radiation of those portions of the composition which are to be removed during development and to permit exposure actinic radiation of those portions which are to remain after development.

The actinic radiation employed in selected such that photopolymerization is accomplished within a reasonable time of from about one second to about 5 minutes. Since free-radical generating addition-polymerization initiators activatable by actinic radiation generally exhibit their maximum sensitivity in the ultraviolet range, the radiation source should furnish an effective amount of this radiation. Both point or broad radiation sources are effective. Such sources include carbon arcs, mercury-vapor arcs, fluorescent lamps with ultraviolet radiation-emitting phosphors, argon glow lamps, electronic flash units and photographic flood lamps. Of these, the mercury-vapor arcs, particularly the sun lamps, are most suitable. In certain circumstances it may be advantageous to expose with visible light, using a photoinitiator sensitive in the visible region of the spectrum, e.g., 9,10-phenanthraquinone. In such cases, the radiation source should furnish an effective dose of visible radiation. Many of the radiation sources listed about furnish the required amount of light energy.

As stated previously, one of the significant advantages of the present invention is that the composition remains flexible upon photopolymerization. This advantage is important in view of the possible uses of the composition and the processing steps commonly encountered. In particular, where flexible substrates are employed, flexing of the laminated, exposed substrates will not cause cracking of the photopolymerized composition. For example, in high volume production, the photopolymerized composition of the present invention can withstand processing performed on a continuous basis wherein the exposed laminate is rolled onto a collection drum prior to further processing. Even where flexible substrates are not employed, the flexible nature of the photopolymerized composition helps to prevent chipping as the laminated,
exposed substrates are handled (e.g., by stacking) and effectively protects the inner surfaces of through
hole connections and other punch outs during the processing of the exposed laminates.

The improved flexibility of the photopolymerized composition of the present invention can be
achieved by employing a more aggressive aqueous alkaline solution than which is used in the developing
step. Thus, for example, while the photopolymerized composition of the present invention is
subjected to etching or plating, it should be apparent that other etching or plating solutions can likewise be used to modify the exposed substrate substantially
without causing deterioration of the resist. Other etching or plating solutions may likewise be used or the composition can be formulated such that it will develop successfully in even more aggressive alkaline solutions.

Suitable alkaline solutions contain from about 0.01 to about 10%, preferably from about 0.5% to about
3% by weight of alkaline agents such as alkali metal hydroxides, e.g., lithium, sodium and potassium
hydroxide; the base-reacting alkali metal salts of weak acids, e.g., lithium, sodium, and potassium
carbonates and bicarbonates; amines having a base-ionization constant greater than about 1 × 10^{-14}, e.g.,
primary amines, such as, benzyl, butyl and allyl amines; secondary amines, e.g., dimethylamine and benzyl
methyl amine; tertiary amines, e.g., trimethylamine, and triethyamine; primary, secondary, and tertiary
hydroxyamines, e.g., propanol, diethanol and triethanol amines, and 2-amino-2-hydroxymethyl-1,3-
propanediol; cyclic amines, e.g., morpholine, piperidine, piperezine, and pyridine; polyamines, such as
hydrizine, ethylene and hexamethylene diamines; the water-soluble basic salts, e.g., the carbonates and
bicarbonates of the above amines; ammonium hydroxide and tetra-substituted ammonium hydroxides,
e.g., tetramethyl-, tetraethyl-, trimethylbenzyl-, and trimethylphenylammonium hydroxides, sulfonium
hydroxides, e.g., trimethylidimethyl-, dimethylbenzylsulfonium hydroxides, and the basic soluble salts
thereof, e.g., the carbonates, bicarbonates and sulfides; alkali metal phosphates and pyrophosphates, e.g.,
sodium and potassium triphosphates and sodium and potassium pyrophosphates; tetra-substituted
(preferably wholly alkyl) phosphonium, arsonium, and stibonium hydroxide, e.g., tetramethyl-
phosphonium hydroxide.

After the development step, the substrate may be recovered, if it is to be used as a printing plate, or may
be subjected to etching or plating if a printed circuit board is to be prepared. Of course, the particular
substrate is initially selected depending on the desired end use.

In the preparation of printed circuit boards, the developed substrate can be passed through a series
of chemical baths to reactivate the exposed metal (e.g., copper). A further advantage of the composition of
the present invention is that the unexposed portions of the composition develop cleanly leaving little or no
residue on the metal surface thereby facilitating reactivation. Moreover, the photopolymerized portions of
the composition are substantially unaffected by the 1—5% by weight solutions of hydrochloric acid,
sulfuric acid, ammonium persulfate, optionally with surfactants, commonly found in the reactivating bath.

Etching or plating is accomplished in those conventional environments known to those of ordinary skill
in the art. That is, while the exposed (i.e., photopolymerized) portions of the composition of the present
invention will not be substantially affected by sequential contact with the developing solution, the
reactivating solution, if used, and a 45° Baume etching solution of ferric chloride, it should be apparent that
other etching or plating solutions can likewise be used to modify the exposed substrate substantially
without causing deterioration of the resist. Other etching or plating solutions are exemplified by solutions
of cupric chloride, ammoniacal etchants, ammonium persulfate, chromic-sulfuric acids, copper sulfate and
solder.
understood that other alkaline solutions, such as those formulated from the alkaline agents disclosed above, can likewise be employed.

The composition of the present invention can also be used as a resist in the process of chemical machining of parts (which are often too small to be acceptably stamped) whereby a metallic sheet or foil is deeply etched or even completely etched through from one side to the other. In the process, the photopolymerizable composition is applied to both sides of the metallic sheet or foil and exposed through identical photographic transparencies as is commonly known in the art. The unexposed portions of the composition are removed in a developing step, as previously described, and the bare surfaces of the metallic sheet or foil are etched as required. Thereafter, the photopolymerized portions of the composition are stripped and the finished product may be recovered and used.

Another important advantage of the present invention pertains to the ease with which the exposed portions of the composition can be stripped from the substrate. This advantage enables the overall processing time to be further reduced which is a decided advantage in the art. By employing the defined acrylate monomer of the present invention in the addition polymerizable material, the stripping time can be reduced by about 10 to about 80%, when compared to the same composition, but not containing the acrylate monomer, under the same stripping conditions.

From the foregoing description of the photopolymerizable composition of the present invention and how it may be formed into sheets or rolls, laminated to a substrate, exposed, developed, subjected to etching or plating and stripped from the substrate, it should be apparent that the photopolymerizable composition of the present invention does not require additional chemical moieties, such as salt groups, in order for it to successfully meet these requirements. Hence, while conventional additives, such as pigments, dyes, adhesion promoters, plasticizers, etc, may be added to enhance the composition of the present invention, the composition consists essentially of the claimed ingredients since it does not require additional chemical moieties, such as salt groups, in order to be employed.

In order to further illustrate the present invention and the advantages thereof, the following specific examples are given, it being understood that these examples are intended only to be illustrative without serving as a limitation on the scope of the present invention.

Example I

The following solutions are coated onto a 25.4 μm (1 mil) thick polyester film, and dried in a current of hot air, provided by a GE—1500 watt fan drier for 20 minutes. The dry thickness of the sensitized layers is about 33 μm (1.3 mils). The dried layers are covered with a 1 mil thick polyethylene film. Formula IA is illustrative of the invention while Formula IB is a control solution.

**Formula IA**

(a) Copolymer of 37% styrene and 63% monobutyl maleate, average mol. wt. 20,000, viscosity of 10% aqueous solution of ammonium salt = 0.15 Pas 38.5 g

(b) Copolymer of 95% vinyl acetate and 5% crotonic acid, average mol. wt. 90,000, viscosity of 8.6% ethyl alcohol solution between 13 and 18 cps 1.5 g

(c) 2,2’-methylene-bis-(4-ethyl-6-tert-butylphenol) 0.06 g

(d) 2-Chlorothioxanthone 0.5 g

(e) Ethyl p-dimethylaminobenzoate 4.0 g

(f) Hecto Blue B dye 0.04 g

(g) Crystal Violet Base 0.015 g

(h) Tripropylene glycol diacrylate 15.0 g

(i) Phenoxydiethoxyethyl acrylate 10.0 g

(j) Methyl ethyl ketone 70.0 g
Formula IB

(a) Copolymer of 37% styrene and 63% monobutyl maleate, average vol. wt. 20,000, viscosity of 10% aqueous solution of ammonium salt = 0.15 Pas

(b) Copolymer of 95% vinyl acetate and 5% crotonic acid, average mol. wt. 90,000, viscosity of 8.6% ethyl alcohol solution between 13 and 18 m Pas

(c) 2,2'-methylene-bus-(4-ethyl-6-tert-butyl phenol) 0.6 g

(d) 2-Chlorothioxanthone 0.5 g

(e) Ethyl p-dimethylaminobenzoate 4.0 g

(f) Hecto Blue B dye 0.04 g

(g) Crystal Violet Base 0.015 g

(h) Tripropylene glycol diacrylate 25.0 g

(i) Methyl ethyl ketone 70.0 g

A piece of copper-clad, epoxy fiberglass board is cleaned by scouring with an abrasive cleaner, swabbing, and thoroughly rinsing in water. It is then given a 20 second dip in a 12% solution of hydrochloric acid, rinsed again with water, and dried with air jets.

The polyethylene cover sheet is removed from a section of the sandwiched photopolymerizable element. The bared resist coating with its polyester support is laminated to the clean copper with the surface of the photopolymerizable layer in contact with the copper surface, using rubber covered rollers operating at a rate of 61 cm per minute at 121°C, with a pressure of 53.6 kg per lineal metre at the nip. The resulting sensitized copper-clad board, protected by the polyester film, is exposed to light through a high-contrast transparency for a period of 30 seconds using a 400 watt, 50 ampere mercury vapor lamp at a distance of 30.5 cm.

The polyester (polyethylene terephthalate) support film is peeled off, and the exposed resist layer is developed by agitating the board in a tray containing a 1% aqueous solution of sodium carbonate monohydrate at 27°C for approximately 1 minute followed by a water rinse. This method of development is satisfactory for the composition of Formula IA as well as the composition of Formula IB. The developed resists have a solid step of 8 and a copper step of 9 on a 21-step scale. The resists reproduce the 0.254 mm lines from the artwork within 2%.

The developed boards are etched in a 45° Baume ferric chloride etchant and the photopolymerized portions of the composition withstand the process without any signs of deterioration or breakdown. The resists are then stripped in an aqueous 3% potassium hydroxide solution at 55°C. The resist made according to Formula IA is stripped off in 27 seconds whereas the resist made according to Formula IB is stripped off in 30 seconds. The etched copper lines in both cases are straight and well-defined.

To measure flexibility, films prepared according to Formulas IA and IB are both laminated to cleaned flexible copper substrates, exposed without artwork to an equivalent of a step 9 on a Stouffer scale of 21, then compared for flexibility using a Gardner MG-1416 Conical Mandrel. The film prepared according to Formula IB shows a crack 48 mm long, whereas the film prepared according to Formula IA shows no crack at all.

Example II

The following solutions are coated onto 25.4 mm thick polyester film, and dried in a current of hot air, provided by a GE-1500 watt fan drier for 20 minutes. The dry thickness of the sensitized layers is about 50.8 μm. The dried layers are covered with a 25.4 μm thick polyethylene film. Formula IIA is illustrative of the invention while Formula IIB is a control solution.
The resist films are laminated, exposed, and developed according to the method described in Example 5.

The surface of the exposed copper is further cleaned by dipping the boards into a 20% ammonium persulfate bath for 30 seconds, washing copiously with water, dipping for 30 seconds in a 20% solution of hydrochloric acid in water, rinsing with water, then drying the boards with jets of air. The cleaned boards are immersed in an acid copper sulfate plating solution containing 19.6 g/l of copper. The plating is carried out at 25°C for 45 minutes using 322 amps per square metre. The boards are taken from the plating tank, water rinsed, and dipped for one minute into a 30% fluoroboric acid solution. The boards are transferred to a hi-throw tin/lead plating bath and plated for 15 minutes. The plating solution contained 16 g/l of tin, 11 g/l of lead and 385 g/l of fluoboric acid. The electric current is 161 amps per square metre. After the plating cycle is completed, the boards are water-rinsed and air-dried. The resists are subjected to a Scotch Tape test to see if any lifting takes place. No lifting is seen for either Formula IIA or IIB. However, when stripped in an aqueous 3% potassium hydroxide solution at 55°C, the resist of Formula IIA is stripped off in 70 seconds, whereas the resist of Formula IIB is stripped off in 165 seconds.
Example III

The following solutions are coated onto a 1 mil thick polyester film, and dried in a current of hot air, provided by a GE-1500 watt fan drier for 20 minutes. The dry thickness of the sensitized layers is about 50.8 \( \mu m \). The dried layers are covered with a 25.4 \( \mu m \) thick polyethylene film. Formula IIIA is illustrative of the invention while Formula IIIB is a control solution.

**Formula IIIA**

(a) Copolymer of 37\% styrene and 63\% monobutyl maleate, average mol. wt. 20,000, viscosity of 10\% aqueous solution of ammonium salt = 0.15 Pas 38.5 g

(b) Copolymer of 95\% vinyl acetate and 5\% crotonic acid, average mol. wt. 90,000, viscosity of 8.6\% ethyl alcohol solution between 13 and 18 mPas 1.5 g

(c) Leuco Crystal Violet dye 0.4 g

(d) Victoria Blue Base dye 0.03 g

(e) Tripropyleneglycol diacrylate 3.0 g

(f) Phenoxydiethoxyethyl acrylate 14.0 g

(g) 2,2'-methylene-bis-(4-ethyl-6-tert-butylphenol) 0.12 g

(h) Benzotriazole 0.1 g

(i) 2,2-Dimethoxy-2-phenylacetophenone 4.0 g

(j) Methyl ethyl ketone 63.0 g

**Formula IIB**

(a) Copolymer of 37\% styrene and 63\% monobutyl maleate, average mol. wt. 20,000, viscosity of 10\% aqueous solution of ammonium salt = 0.15 Pas 38.5 g

(b) Copolymer of 95\% vinyl acetate and 5\% crotonic acid, average mol. wt. 90,000, viscosity of 8.6\% ethyl alcohol solution between 13 and 18 mPas 1.5 g

(c) Leuco Crystal Violet dye 0.4 g

(d) Victoria Blue Base FB dye 0.03 g

(e) Tripropyleneglycol diacrylate 8.0 g

(f) Diethoxylated Bisphenol-A diacrylate 14.0 g

(g) 2,2'-methylene-bis-(4-ethyl-6-tert-butylphenol) 0.12 g

(h) Benzotriazole 0.1 g

(i) 2,2-Dimethoxy-2-phenylacetophenone 4.0 g

(j) Methyl ethyl ketone 63.0 g

As described in Example II, 50.8 \( \mu m \) thick films are prepared and evaluated as plating resists. Both Formulas IIIA and IIB performed satisfactorily exhibiting no lifting as shown by the Scotch Tape test and by the absence of underplating. When stripped under the conditions described in Example II, the resist of Formula IIIA is stripped off in 57 seconds, whereas the resist of Formula IIB is stripped off in 70 seconds.

Example IV

The following solutions are coated onto a 25.4 \( \mu m \) thick polyester film, and dried in a current of hot air, provided by a GE-1500 watt fan drier for 20 minutes. The dry thickness of the sensitized layers is about 50.8 \( \mu m \). The dried layers are covered with a 25.4 \( \mu m \) thick polyethylene film. Formula IVA is illustrative of the invention while Formula IVB is a control solution.
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Formula IVA
(a) Copolymer of 37% styrene and 63% monobutyl maleate, average mol. wt.
20,000, viscosity of 10% aqueous solution of ammonium salt = 0.15 Pas 38.5 g
5
(b) Copolymer of 95% vinyl acetate and 5% crotonic acid, average mol. wt.
90,000, viscosity of 8.6% ethyl alcohol solution between 13 and 18 mPas 1.5 g
(c) Leuco Crystal Violet dye 0.4 g
(d) Victoria Blue Base FB 0.03 g
(e) Trimethylolpropane triacrylate 10.5 g
(f) Phenoxydiethoxyethyl acrylate 10.5 g
(g) 2,2'-methylene-bis-(4-ethyl-6-tert-butylphenol) 0.12 g
(h) Benzotriazole 0.1 g
(i) 2,2-Dimethoxy-2-phenylacetonophenone 4.0 g
(j) Methyl ethyl ketone 63.0 g

Formula IVB
(a) Copolymer of 37% styrene and 63% monobutyl maleate, average mol. wt.
20,000, viscosity of 10% aqueous solution of ammonium salt = 0.15 Pas 38.5 g
5
(b) Copolymer of 95% vinyl acetate and 5% crotonic acid, average mol. wt.
90,000, viscosity of 8.6% ethyl alcohol solution between 13 and 18 mPas 1.5 g
(c) Leuco Crystal Violet dye 0.4 g
(d) Victoria Blue Base FB 0.03 g
(e) Tripropyleneglycol diacylate 8.0 g
(f) Triethyleneglycol diacylate 10.5 g
(g) 2,2'-methylene-bis-(4-ethyl-6-tert-butylphenol) 0.12 g
(h) Benzotriazole 0.1 g
(i) 2,2-Dimethoxy-2-phenylacetonophenone 4.0 g
(j) Methyl ethyl ketone 63.0 g

The two formulas are tested as plating resists according to the methods described in Examples II and III. Both films performed satisfactorily in the plating cycle, and are then stripped under the same conditions as described in Example II. The resist of Formula IVA is stripped off in 50 seconds and the resist of Formula IVB is stripped off in 128 seconds.

Example V
The following solutions are coated onto 25.4 μm thick polyester film, and dried in a current of hot air, provided by a GE-1500 watt fan drier, for 20 minutes. The dry thickness of the sensitized layers is about 50.8 μm. The dried layers are covered with 25.4 μm thick polyethylene film. Formula VA is illustrative of the invention, while Formula VB is a control solution.
The two formulas performed satisfactorily when evaluated as plating resists as in Example II. When subjected to stripping conditions as described in Example II, the resist of Formula VA is stripped off in 54 seconds, whereas the resist of Formula VB is stripped off in 100 seconds. Separate samples of films prepared from Formulas VA and VB are subjected to the flexibility test as described in Example I. The length of the crack with Formula VA is 13.4 cm, thus showing the superior flexibility of Formula VA.
Claims

1. A photopolymerizable composition capable of being formed into a storable sheet or roll on a peelable support which on a 100 parts by weight basis comprises:

A. from 10 to 60 parts by weight of an addition polymerizable material including one or more non-gaseous compounds containing at least two terminal ethylenic groups and having a boiling point above 100°C;

B. from 0.001 to 20 parts by weight of a photo initiated free radical generating addition polymerization initiating system;

C. from 0.001 to 5 parts by weight of a thermal addition polymerization inhibitor; and

D. from 40 to 90 parts by weight of a preformed macro-molecular polymeric binding agent which is a polymer of:

i) a first monomeric material which contains one or more non acidic compounds having the general formula

\[
\begin{align*}
R-C=CH_2
\end{align*}
\]

or

\[
\begin{align*}
CH_2-C-Y
\end{align*}
\]

wherein \( R \) is hydrogen, an alkyl group having from 1 to 6 carbon atoms or a halo group and wherein when \( X \) is hydrogen \( Y \) is OOCR, OR, OCR, COOR, CN, CH=CH2

\[
\begin{align*}
O
\end{align*}
\]

\[
\begin{align*}
\text{CNR}_3 \text{R}_4
\end{align*}
\]

or CI, when \( X \) is methyl, \( Y \) is COOR, CN, CH=CH2 or

\[
\begin{align*}
O
\end{align*}
\]

\[
\begin{align*}
\text{CNR}_3 \text{R}_4
\end{align*}
\]

and when \( X \) is chlorine, \( Y \) is CI and wherein \( \text{R}_1 \) is an alkyl group having from 1 to 12 carbon atoms, a phenyl group or a benzyl group and \( \text{R}_3 \) and \( \text{R}_4 \) are hydrogen, an alkyl group having from 1 to 12 carbon atoms or a benzyl group; and

ii) a second monomeric material which consists essentially of one or more alpha, beta ethylenically unsaturated carboxylic acid- or anhydride-containing monomers having from 3 to 15 carbon atoms, the ratio of the first monomeric material to the second monomeric material being sufficient to render substantially all of the binding agent soluble in a substantially wholly aqueous solution containing 1% by weight of sodium carbonate, characterised in that component A comprises:

i) from 5 to 50 parts by weight of an acrylate having a boiling point greater than 175°C and having the formula

\[
\begin{align*}
\text{R}_1
\end{align*}
\]

\[
\begin{align*}
\text{H}_2\text{C}=-\text{C}-(\text{OC}_m\text{H}_2\text{m})_n\text{ OR}_2
\end{align*}
\]

wherein \( m \) is from 1 to 4, \( n \) is from 1 to 12, \( \text{R}_1 \) is selected from H, CH3 and mixtures thereof and \( \text{R}_2 \) is selected from unsubstituted phenyl, substituted phenyl, unsubstituted naphthyl, substituted naphthyl, branched or unbranched, substituted or unsubstituted alkyl having 1 to 12 carbon atoms, substituted or unsubstituted cycloalkyl having 5 or 6 carbon atoms in the ring and mixtures thereof and

ii) 5 to 50 parts by weight of said non gaseous compounds containing two or more terminal ethylenic groups.

2. The photopolymerizable composition of claim 1 wherein \( n \) is from 2 to 8 and \( \text{R}_2 \) is substituted or unsubstituted phenyl.

3. The photopolymerizable composition of claim 2 wherein \( m \) is 2 or 3, \( n \) is 2 or 3, \( \text{R}_1 \) is H and \( \text{R}_2 \) is unsubstituted phenyl.

4. The photopolymerizable composition as claimed in any preceding claim, wherein the non-gaseous compounds containing at least two terminal ethylenic groups and having a boiling point above 100°C are selected from unsaturated ester of a polyl, an unsaturated amide, a vinyl ester and mixtures thereof.
5. The photopolymerizable composition of claim 4, wherein the non-gaseous compounds are selected from trimethylolpropane triacrylate, tetraethyleneglycol diacrylate, tripropyleneglycol diacrylate and mixtures thereof.

6. The photopolymerizable composition of claim 4 wherein the non-gaseous compounds comprise tripropyleneglycol diacrylate.

7. The photopolymerizable composition of claim 4 wherein the non-gaseous compounds comprise pentaerythritol acrylates.

8. The photopolymerizable composition as claimed in any preceding claim wherein the binding agent has the first monomeric material having the general formula

\[ R-C = CH_2 \]

9. The photopolymerizable composition of claim 8 wherein the binding agent is a copolymer of styrene and monobutyl maleate in a ratio of from 35:65 to 70:30.

10. The photopolymerizable composition as claimed in any one of claims 1 to 7 wherein the binding agent is a copolymer of vinyl acetate and crotonic acid in a ratio of from 70:30 to 95:5.

11. The photopolymerizable composition as claimed in any one of claims 1 to 7 wherein the binding agent is a mixture of a copolymer of styrene and monobutyl maleate in a ratio of from 35:65 to 70:30 with a copolymer of vinyl acetate and crotonic acid in a ratio of from 70:30 to 95:5.

12. The photopolymerizable composition as claimed in any one of claims 1 to 7 wherein the binding agent is a copolymer of an acrylate-type monomer and one or more alpha, beta-ethylenically unsaturated carboxylic acid- or anhydride-containing monomers having from 3 to 15 carbon atoms.

13. The photopolymerizable composition as claimed in any one of claims 1 to 7 wherein the binding agent is a copolymer of

i) an acrylate-type monomer and

ii) one or more alpha, beta-ethylenically unsaturated carboxylic acid- or anhydride-containing monomers having from 3 to 15 carbon atoms.

14. A sheet or roll of a photopolymerizable composition on a flexible film support, wherein said photopolymerizable composition is as defined in any preceding claim and is peelable from the support.

15. The sheet or roll of claim 14 wherein the acrylate is selected from substituted and unsubstituted phenoxypolyethoxyethyl acrylates, substituted and unsubstituted phenoxypolypropoxypropyl acrylates and mixtures thereof.

16. A substrate coated with a photopolymerizable composition, wherein said photopolymerizable composition is defined in any one of claims 1 to 14.

17. The coated substrate of claim 16 wherein the substrate is comprised of a metal.

18. The coated substrate of claim 16 wherein the substance is comprised of a metal.

19. The coated substrate of claim 15 wherein the metal is selected from steel, aluminium, copper, zinc and magnesium.

20. The coated substrate of claim 15 wherein the metal is selected from steel, aluminium, copper, zinc and magnesium.

21. The coated substrate of claim 18 wherein the substrate is comprised of plastics.

22. The coated substrate of claim 19 wherein the substrate is a copper clad laminate.

23. The coated substrate of claim 20 wherein the substrate is a copper clad, epoxy-fiberglass laminate.

24. A process for preparing a printed circuit board comprising:

A. laminating onto a printed circuit board precursor a photopolymerizable composition as claimed in any one of claims 1 to 14;

B. exposing selected portions of the photopolymerizable composition to actinic radiation sufficient to obtain photopolymerization of said exposed portions;

C. removing the unexposed portions of the composition to bare portions of the circuit board precursor by contact with a substantially wholly aqueous solution having an alkalinity at least as great as a 1% by weight solution of sodium carbonate;
D. modifying the bared portions of the circuit board precursor to form a selective electrically conductive pattern; and
E. stripping the exposed selected portions of the composition from the circuit board by contact with a substantially wholly aqueous solution having an alkalinity at least as great as a 1% by weight solution of sodium hydroxide.

25. The process of claim 24 wherein the printed circuit board precursor is a copper clad, epoxy fiberglass laminate.
26. The process of claim 24 wherein the printed circuit board precursor is a flexible, copper clad laminate.
27. The process as claimed in any one of claims 24 to 26 wherein the printed circuit board precursor has a metallic material on both sides and the photopolymerizable composition is laminated to both sides on top of said metallic material.
28. The process as claimed in any one of claims 24 to 27 wherein the photopolymerizable composition is laminated to the printed circuit board precursor in the form of a sheet or a portion of a roll on a film support.
29. The process as claimed in any one of claims 24 to 28 wherein the bared portions of the circuit board precursor are modified by etching.
30. The process as claimed in any one of claims 24 to 28 wherein the bared portions of the circuit board precursor are modified by plating.

31. A process for chemical machining comprising:
A. laminating onto both sides of a metallic sheet or foil a polymerizable composition as claimed in any one of claims 1 to 14;
B. exposing selected portions of the photopolymerizable composition to actinic radiation sufficient to obtain photopolymerization of said exposed portions;
C. removing the unexposed portions of the composition to bare portions of the sheet or foil by contact with a substantially wholly aqueous solution having an alkalinity at least as great as a 1% by weight solution of sodium carbonate;
D. etching the bared portions of the sheet or foil to form the desired pattern; and
E. stripping the exposed selected portions of the composition from the etched sheet or foil by contact with a substantially wholly aqueous solution having an alkalinity at least as great as a 1% by weight solution of sodium hydroxide.

Patentansprüche

1. Photopolymerisierbare Zusammensetzung, die zu einem lagerfähigen Bogen oder einer lagerfähigen Rolle auf einem abziehbaren Träger verarbeitet werden kann und die auf der Grundlage von 100 Gewichtsteilen
   A. 10 bis 60 Gewichtsteile eines additionspolymerisierbaren Materials einschließlich einer oder mehrerer nichtgasförmiger Verbindungen, die wenigstens zwei endständige ethylenische Gruppen enthalten und einen Siedepunkt oberhalb 100°C haben,
   B. 0,001 bis 20 Gewichtsteile eines durch Photoinitiierung freie Radikale erzeugenden, die Additionspolymerisation initiierenden Systems,
   C. 0,001 bis 5 Gewichtsteile eines Inhibitors für thermische Additionspolymerisation und
   D. 40 bis 90 Gewichtsteile eines vorgeformten makromolekularen polymeren Bindemittels umfaßt, das ein Polymer
   i) eines ersten Monomermaterials, das eine oder mehrere nichtsäure Verbindungen der allgemeinen Formel

   \[
   \begin{align*}
   R - C &= CH_2 \\
   CH_2 &= C - Y \\
   X &= \ \text{oder} \\
   O &= \\
   CNR_3R_4 &= 
   \end{align*}
   \]

   enthält, worin R Wasserstoff, eine Alkylgruppe mit 1 bis 6 Kohlenstoffatomen oder eine Halogengruppe ist und worin, wenn X Wasserstoff ist, Y OOCR₁, OR₁, OOCR₁, COOR₁, CN, CH=CH₂, oder

   \[
   \begin{align*}
   O &= \\
   CNR_3R_4 &= 
   \end{align*}
   \]

   ist, und, wenn X Chlor ist, Y Cl ist, und worin R₁ eine Alkylgruppe mit 1 bis 12 Kohlenstoffatomen, eine Phenylgruppe oder eine Benzylgruppe ist und R₂ und R₄ Wasserstoff, eine Alkylgruppe mit 1 bis 12 Kohlenstoffatomen oder eine Benzylgruppe sind, und
ii) eines zweiten Monomermaterials, das im wesentlichen aus einem oder mehreren alpha-, beta-
ethylenisch ungesättigten carbonsäure- oder anhydridhaltigen Monomeren mit 3 bis 15
Kohlenstoffatomen, wobei das Verhältnis des ersten Monomermaterials zu dem zweiten Monomermaterial
ausreicht, um im wesentlichen das gesamte Bindemittel in einer im wesentlichen vollständig wäßrigen
Lösung mit einem Gehalt von 1 Gew.-% Natriumcarbonat löslich zu machen, ist, dadurch gekennzeichnet,
däß die Komponente A
i) 5 bis 50 Gewichtsteile eines Acrylates mit einem Siedepunkt größer als 175°C und mit der Formel

\[
\begin{align*}
H_2C=\overset{\delta}C-\overset{\delta}C-(OC_mH_{2m})_nOR_2
\end{align*}
\]

worin m 1 bis 4 ist, n 1 bis 12 ist, R₁ unter H, CH₃ und Gemischen hiervon ausgewählt ist und R₂ unter
unsubstituiertem Phenyl, substituiertem Phenyl, unsubstituiertem Naphthyl, substituiertem Naphthyl,
verzweigtem oder unverzweigtem, substituiertem oder unsubstituiertem Alkyl mit 1 bis 12
Kohlenstoffatomen, substituiertem oder unsubstituiertem Cycloalyl mit 5 oder 6 Kohlenstoffatomen im
Ring sowie Gemischen hiervon ausgewählt ist, und

ii) 5 bis 50 Gewichtsteile der nichtgasförmigen Verbindungen, die zwei oder mehr endständige
ethylenische Gruppe enthalten, umfaßt.

2. Photopolymerisierbare Zusammensetzung nach Anspruch 1, worin n 2 bis 8 ist und R₂ substituiertes
oder unsubstituiertes Phenyl ist.

3. Photopolymerisierbare Zusammensetzung nach Anspruch 2, worin m 2 oder 3 ist, n 2 oder 3 ist, R₁ H
ist und R₂ unsubstituiertes Phenyl ist.

4. Photopolymerisierbare Zusammensetzung nach einem der vorausgehenden Ansprüche, worin die
nichtgasförmigen Verbindungen, die wenigstens zwei endständige ethylenische Gruppen enthalten und
einen Siedepunkt oberhalb 100°C haben, unter ungesättigtem Ester eines Polyols, ungesättigtem Amid,
Vinyester und Gemischen hiervon ausgewählt sind.

5. Photopolymerisierbare Zusammensetzung nach Anspruch 4, worin die nichtgasförmigen
Verbindungen unter Trimethylolpropantriacylat, Tetraethylenglycoldiacrylat, Tripropylenglycoldiacrylat
und Gemischen hiervon ausgewählt sind.

6. Photopolymerisierbare Zusammensetzung nach Anspruch 4, worin die nichtgasförmigen
Verbindungen Tripropylenglycoldiacrylat umfassen.

7. Photopolymerisierbare Zusammensetzung nach Anspruch 4, worin die nichtgasförmigen
Verbindungen Pentaerythritacrylate umfassen.

8. Photopolymerisierbare Zusammensetzung nach einem der vorausgehenden Ansprüche, worin das
Bindemittel das erste Monomermaterial der allgemeinen Formel

\[
R-C=CH_2
\]

hat.

9. Photopolymerisierbare Zusammensetzung nach Anspruch 8, worin das Bindemittel ein Copolymer
von Styrol und Monobutylmaleat im Verhältnis von 35:65 bis 70:30 ist.

10. Photopolymerisierbare Zusammensetzung nach einem der Ansprüche 1 bis 7, worin das
Bindemittel ein Copolymer von Vinylacetat und Crotonsäure in einem Verhältnis von 70:30 bis 95:5 ist.

11. Photopolymerisierbare Zusammensetzung nach einem der Ansprüche 1 bis 7, worin das
Bindemittel ein Gemisch eines Copolymers von Styrol und Monobutylmaleat in einem Verhältnis von 35:65
bis 70:30 mit einem Copolymer von Vinylacetat und Crotonsäure in einem Verhältnis von 70:30 bis 95:5 ist.

12. Photopolymerisierbare Zusammensetzung nach einem der Ansprüche 1 bis 7, worin das
Bindemittel ein Copolymer eines Monomers vom Acrylattyp und eines oder mehrerer alpha-, beta-
ethylenisch ungesättigten carbonsäure- oder anhydridhaltiger Monomere mit 3 bis 15 Kohlenstoffatomen
ist.

13. Photopolymerisierbare Zusammensetzung nach einem der Ansprüche 1 bis 7, worin das
Bindemittel ein Copolymer von

\[
R-C=CH_2
\]

ist.
50 Schaltungsplattenvorläufer durch Atzen modifiziert werden.
55 beiden Seiten eines Metallbogens oder einer Metallfolie, auf die Zusammensetzung auf beide Seiten auf das metallische Material aufgebracht wird. 

15 selektiven elektrisch leitenden Musters und

22 Verfahren zur Herstellung einer gedruckten Schaltungsplatte unter

24 Verfahren zur Herstellung einer gedruckten Schaltungsplatte unter

25 Verfahren zur Herstellung einer gedruckten Schaltungsplatte unter

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Revendications

1. Composition photopolymérisable capable d’être façonnée en une feuille ou un rouleau propres à la conservation sur un support pelable qui, sur base de 100 parties en poids, comprend:
   a. 10 à 60 parties en poids d’une matière photopolymérisable par addition comprenant un ou plusieurs composés non gazeux contenant au moins deux radicaux éthyléniques terminaux et ayant un point d’ébullition supérieur à 100°C;
   B. 0,001 à 20 parties en poids d’un système initiateur de polymérisation par addition engendrant des radicaux libres photo-initiés;
   C. 0,001 à 5 parties en poids d’un inhibiteur de polymérisation par addition thermique; et
   D. 40 à 90 parties en poids d’un agent liant polymère macromoléculaire préformé qui est un polymère de
   i) une première matière monomère qui contient un ou plusieurs composés non acides de formule générale
   ![Structure de la première matière monomère](image)
   ou
   ![Structure de la deuxième matière monomère](image)
   où R est un atome d’hydrogène, un radical alcoyle de 1 à 6 atomes de carbone ou un radical halo et où lorsque X est un atome d’hydrogène, Y est OOCR1, OR1, OCR1, COOR1, CN, CH=CH2, O=CNR3R4 ou Cl, lorsque X est un radical méthyle, Y est COOR1, CN, CH=CH2 ou O=CNR3R4 et
   lorsque X est un atome de chlore, Y est Cl et où R1 est un radical alcoyle de 1 à 12 atomes de carbone, un radical phényle ou un radical benzyle et R2 et R3 sont un atome d’hydrogène, un radical alcoyle de 1 à 12 atomes de carbone ou un radical benzyle; et
   ii) une seconde matière monomère qui consiste essentiellement en un ou plusieurs monomères alpha-, bêta-éthyléniquement insaturés contenant la fonction acide ou anhydride carboxylique de 3 à 15 atomes de carbone, le rapport de la première matière monomère à la seconde matière monomère étant suffisant pour rendre sensiblement l’ensemble de l’agent liant soluble dans une solution sensiblement entièrement aqueuse contenant 1% en poids de carbonate de sodium, caractérisée en ce que le composant A comprend:
   i) 5 à 50 parties en poids d’un acrylate ayant un point d’ébullition supérieur à 175°C et répondant à la formule
   ![Structure de l’acrylate](image)
   où m est de 1 à 4, n est de 1 à 12, R1 est choisi entre H, CH3 et leurs mélanges et R2 est choisi parmi les radicaux phényle non substitué, phényle substitué, naphtyle non substitué, naphtyle substitué, alcoyle de 1 à 12 atomes de carbone ramifié ou non ramifié substitué ou non substitué, cycloalcoyle de 5 ou 6 atomes de carbone dans le cycle substitué ou non substitué et leurs mélanges, et
   ii) 5 à 50 parties en poids des dits composés non gazeux contenant deux ou plusieurs radicaux éthyléniques terminaux.

2. Composition photopolymérisable suivant la revendication 1, dans laquelle n est de 2 à 8 et R2 est un radical phényle substitué non substitué.

3. Composition photopolymérisable suivant la revendication 2, dans laquelle m est 2 ou 3, n est 2 ou 3, R1 est H et R2 est un radical phényle non substitué.

4. Composition photopolymérisable suivant l’une quelconque des revendications précédentes, dans laquelle les composés non gazeux contenant au moins deux radicaux éthyléniques terminaux et ayant un point d’ébullition supérieur à 100°C sont choisis parmi un ester insaturé d’un polyl, un amide insaturé, un ester vinylique et leurs mélanges.

5. Composition photopolymérisable suivant la revendication 4, dans laquelle les composés non gazeux sont choisis parmi le triacrylate de triméthylolpropane, le diacrylate de tétraéthyléneglycol, le diacrylate de tripropyléneglycol et leurs mélanges.

6. Composition photopolymérisable suivant la revendication 4, dans laquelle les com posés non gazeux comprennent du diacrylate de tripropyléneglycol.

7. Composition photopolymérisable suivant la revendication 4, dans laquelle les composés non gazeux comprennent des acrylates de pentaérythritol.

8. Composition photopolymérisable suivant l’une quelconque des revendications précédentes, dans laquelle l’agent liant comprend la première matière monomère de formule générale
9. Composition photopolymérisable suivant la revendication 8, dans laquelle l'agent liant est un copolymère de styrène et de maléate de monobutyle dans un rapport de 35:65 à 70:30.

10. Composition photopolymérisable suivant l'une quelconque des revendications 1 à 7, dans laquelle l'agent liant est un copolymère d'acétate de vinyle et d'acide crotonique dans un rapport de 70:30 à 95:5.

11. Composition photopolymérisable suivant l'une quelconque des revendications 1 à 7, dans laquelle l'agent liant est un mélange d'un copolymère de styrène et de maléate de monobutyle dans un rapport de 35:65 à 70:30 avec un copolymère d'acétate de vinyle et d'acide crotonique dans un rapport de 70:30 à 95:5.

12. Composition photopolymérisable suivant l'une quelconque des revendications 1 à 7, dans laquelle l'agent liant est un copolymère d'un monomère du type acrylique et d'un ou plusieurs monomères alpha, beta-éthyléniquement insaturés contenant la fonction acide ou anhydride carboxylique de 3 à 15 atomes de carbone.

13. Composition photopolymérisable suivant l'une quelconque des revendications 1 à 7, dans laquelle l'agent liant est un copolymère

\[ R-C=CH_2 \]

i) d'un monomère du type acrylique et

ii) d'un ou plusieurs monomères alpha, beta-éthyléniquement insaturés contenant la fonction acide ou anhydride carboxylique de 3 à 15 atomes de carbone.

14. Composition photopolymérisable suivant la revendication 13, dans laquelle le monomère du type acrylique est choisi parmi un acrylate d'alcoyle, un méthacrylate d'alcoyle, un acrylate d'hydroxyalcoyle, un méthacrylate d'hydroxyalcoyle et leurs mélanges dont les radicaux alcoyle comptent 1 à 12 atomes de carbone et dont les radicaux hydroxyalcoyle comptent 2 à 12 atomes de carbone.

15. Feuille ou rouleau d'une composition photopolymérisable sur un film de support flexible, dont la composition photopolymérisable est telle que définie dans l'une quelconque des revendications précédentes et est pelable à partir du support.

16. Feuille ou rouleau suivant la revendication 15, dont l'acrylate est choisi parmi les acrylates de phénoxypropyléthoxyéthyle substitués et non substitués, les acrylates de phénoxypropoxypropyle substitués et non substitués et leurs mélanges.

17. Feuille ou rouleau suivant la revendication 16, dont l'acrylate est un acrylate de phénoxypropyléthoxyéthyle non substitué.

18. Substrat revêtu d'une composition photopolymérisable, dans lequel la composition photopolymérisable est telle que définie dans l'une quelconque des revendications 1 à 14.

19. Substrat revêtu suivant la revendication 18, dans lequel le substrat est formé d'un métal.

20. Substrat revêtu suivant la revendication 19, dans lequel le métal est choisi entre l'acier, l'aluminium, le cuivre, le zinc et le magnésium.

21. Substrat revêtu suivant la revendication 18, dans lequel le substrat est formé de matière plastique.

22. Substrat revêtu suivant la revendication 18, dans lequel le substrat est un stratifié doublé de cuivre.

23. Substrat revêtu suivant la revendication 22, dans lequel le substrat est un stratifié fibre de verre-époxyde doublé de cuivre.

24. Procédé de préparation d'une plaque à circuit imprimé, qui comprend:

A. la stratification, sur un précurseur de plaque à circuit imprimé, d'une composition photopolymérisable suivant l'une quelconque des revendications 1 à 14;
B. l'exposition de parties sélectionnées de la composition photopolymérisable a un rayonnement actinique suffisant pour provoquer la photopolymérisation des parties exposées;
C. l'élimination des parties non exposées de la composition pour dénuder des parties du précurseur de plaque à circuit imprimé par contact avec une solution sensiblement complètement aqueuse ayant une alcalinité au moins aussi forte qu'une solution à 1% en poids de carbonate de sodium;
D. la modification des parties dénudées du précurseur de plaque à circuit imprimé pour y former un motif sélectif conducteur de l'électricité, et
E. le dépouillement des parties sélectionnées exposées de la composition de la plaque à circuit imprimé par contact avec une solution sensiblement complètement aqueuse ayant une alcalinité au moins aussi forte qu'une solution à 1% en poids d'hydroxyde de sodium.
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25. Procédé suivant la revendication 24, dans lequel le précurseur de plaque à circuit imprimé est un stratifié fibre de verre-époxyde doublé de cuivre.

26. Procédé suivant la revendication 24, dans lequel le précurseur de plaque à circuit imprimé est un stratifié flexible doublé de cuivre.

27. Procédé suivant l’une quelconque des revendications 24 à 26, dans lequel le précurseur de plaque à circuit imprimé porte une matière métallique sur ses deux faces et la composition photopolymérisable est stratifiée sur les deux faces par-dessus la matière métallique.

28. Procédé suivant l’une quelconque des revendications 24 à 27, dans lequel la composition photopolymérisable est stratifiée sur le précurseur de plaque à circuit imprimé sous la forme d’une feuille ou d’une partie d’un rouleau sur un film de support.

29. Procédé suivant l’une quelconque des revendications 24 à 28, dans lequel les parties dénudées du précurseur de plaque à circuit imprimé sont modifiées par morsure.

30. Procédé suivant l’une quelconque des revendications 24 à 28, dans lequel les parties dénudées du précurseur de plaque à circuit imprimé sont modifiées par dépôt de métal.

31. Procédé d’usinage chimique, qui comprend

A. la stratification, sur les deux faces d’une feuille ou d’un clinquant métalliques, d’une composition photopolymérisable suivant l’une quelconque des revendications 1 à 14;

B. l’exposition de parties sélectionnées de la composition photopolymérisable à un rayonnement actinique suffisant pour provoquer la photopolymérisation des parties exposées;

C. l’élimination des parties non exposées de la composition pour dénuder des parties de la feuille ou du clinquant par contact avec une solution sensiblement complètement aqueuse ayant une alcalinité au moins aussi forte qu’une solution à 1% en poids de carbonate de sodium;

D. la morsure des parties dénudées de la feuille ou du clinquant pour y former le motif désiré;

E. le dépouillement des parties sélectionnées exposées de la composition de la feuille ou du clinquant ayant subi la morsure par contact avec une solution sensiblement complètement aqueuse ayant une alcalinité au moins aussi forte qu’une solution à 1% en poids d’hydroxyde de sodium.